

4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 573

[Docket No. FDA-2014-F-1184]

Zinpro Corp.; Filing of Food Additive Petition (Animal Use)

AGENCY: Food and Drug Administration, HHS.

ACTION: Notification; petition for rulemaking.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing that Zinpro Corp. has filed a petition proposing that the food additive regulations be amended to provide for the safe use of silicon dioxide as an anticaking agent for the use with zinc-L-selenomethionine as a feed component.

DATES: The food additive petition was filed on July 29, 2014.

ADDRESSES: For access to the docket, go to https://www.regulations.gov and insert the docket number, found in brackets in the heading of this document, into the "Search" box and follow the prompts; and/or go to the Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Carissa Doody, Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855, 240-402-6283, Carissa.doody@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the Federal Food, Drug, and Cosmetic Act (section 409(b)(5) (21 U.S.C. 348(b)(5))), notice is given that a food additive petition (FAP 2285) has been filed by Zinpro Corp., 10400 Viking Dr., suite 240, Eden Prairie, MN 55344.

The petition proposes to amend Title 21 of the Code of Federal Regulations (CFR) in part 573

(21 CFR part 573) Food Additives Permitted in Feed and Drinking Water of Animals to provide

for the safe use of silicon dioxide as an anticaking agent for the use with zinc-L-

selenomethionine as a feed component. In an earlier notice of petition (79 FR 49465, August 21,

2014), the use of silicon dioxide was omitted.

The petitioner has claimed that this action is categorically excluded under 21 CFR

25.32(r) because it is of a type that does not individually or cumulatively have a significant effect

on the human environment. In addition, the petitioner has stated that, to their knowledge, no

extraordinary circumstances exist. If FDA determines a categorical exclusion applies, neither an

environmental assessment nor an environmental impact statement is required. If FDA

determines a categorical exclusion does not apply, we will request an environmental assessment

and make it available for public inspection.

Dated: October 25, 2018.

Leslie Kux,

Associate Commissioner for Policy.

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